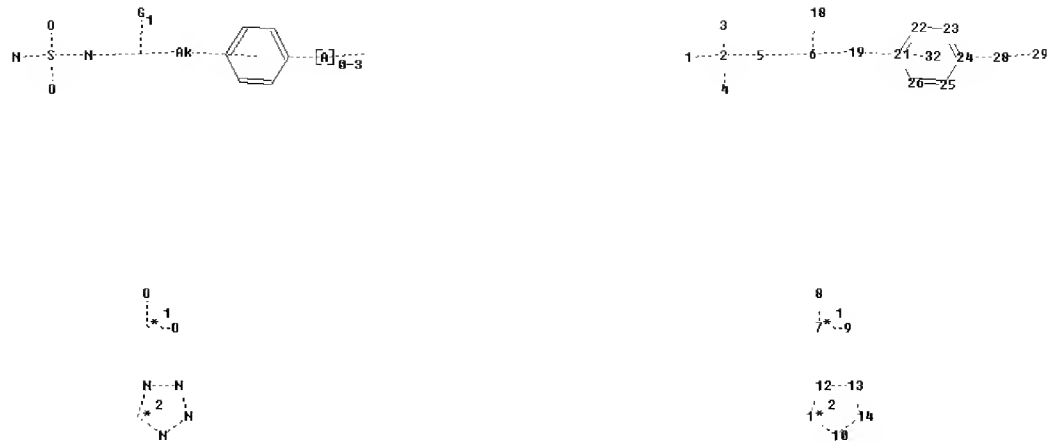


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chain nodes :

2 3 4 5 6 7 8 9 18 19 28

ring nodes :

10 11 12 13 14 21 22 23 24 25 26 29

ring/chain nodes :

1

chain bonds :

1-2 2-3 2-4 2-5 5-6 6-18 6-19 7-8 7-9 24-28 28-29

ring bonds :

10-11 10-14 11-12 12-13 13-14 21-26 21-22 22-23 23-24 24-25 25-26

exact/norm bonds :

1-2 2-3 2-4 2-5 5-6 6-18 6-19 7-8 7-9 10-11 10-14 11-12 12-13 13-14
24-28 28-29

normalized bonds :

21-26 21-22 22-23 23-24 24-25 25-26

isolated ring systems :

containing 10 : 21 :

G1:[*1], [*2]

Connectivity :

19:2 E exact RC ring/chain

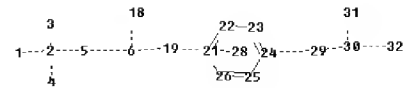
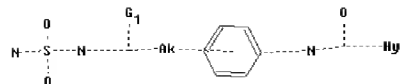
Match level :

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 18:CLASS 19:CLASS 21:Atom 22:Atom
23:Atom 24:Atom
25:Atom 26:Atom 28:CLASS 29:Atom 32:Atom

L1 STRUCTURE UPLOADED

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chain nodes :
 1 2 3 4 5 6 7 8 9 18 19 29 30 31 32
 ring nodes :
 10 11 12 13 14 21 22 23 24 25 26
 chain bonds :
 1-2 2-3 2-4 2-5 5-6 6-18 6-19 7-8 7-9 24-29 29-30 30-31 30-32
 ring bonds :
 10-11 10-14 11-12 12-13 13-14 21-26 21-22 22-23 23-24 24-25 25-26
 exact/norm bonds :
 1-2 2-3 2-4 2-5 5-6 6-18 6-19 7-8 7-9 10-11 10-14 11-12 12-13 13-14
 24-29 29-30 30-31 30-32
 normalized bonds :
 21-26 21-22 22-23 23-24 24-25 25-26
 isolated ring systems :
 containing 10 : 21 :

G1:[*1], [*2]

Connectivity :
 19:2 E exact RC ring/chain
 Match level :
 1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 18:CLASS 19:CLASS 21:Atom 22:Atom
 23:Atom 24:Atom
 25:Atom 26:Atom 28:Atom 29:CLASS 30:CLASS 31:CLASS 32:Atom
 Generic attributes :
 32:
 Saturation : Unsaturated
 Number of Carbon Atoms : less than 7
 Number of Hetero Atoms : Exactly 1
 Type of Ring System : Monocyclic

Element Count :

Node 32: Limited

C,C5
N,N1

=> d his

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L1 STRUCTURE UPLOADED
L2 STRUCTURE UPLOADED
L3 81 S L1 SSS FULL
L4 35 S L2 SSS FULL SUB=L3

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L5 2 S L4
L6 2 S US2001-555286/APPS
L7 2 S L5 AND L6
L9 3 S L3
L10 1 S L9 NOT L6

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L7 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN
AN 2005:1107613 CAPLUS Full-text

DN 143:326627

TI Preparation of N-(2-phenylethyl)sulfamide derivatives as α 4 integrin antagonists for treatment of inflammatory and immune disorders

IN Jimenez Mayorga, Juan Miguel; Vidal Gispert, Laura; Warrelow, Graham

PA Almirall Prodesfarma, S.A., Spain

SO Span., 41 pp.

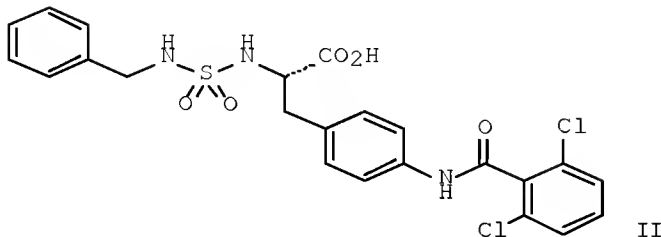
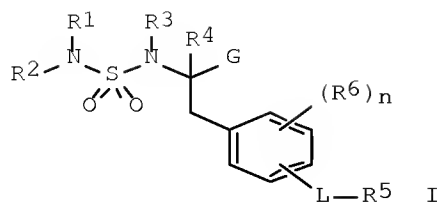
CODEN: SPXXAD

DT Patent

LA Spanish

FAN.CNT 2

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PI	ES 2219177	A1	20041116	ES 2003-1004	20030505
	ES 2219177	B1	20060216		
	WO 2004099126	A1	20041118	WO 2004-EP4670	20040503
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		RW:	BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG		
	EP 1622867	A1	20060208	EP 2004-730833	20040503
	EP 1622867	B1	20070919		
		R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR		
	CN 1816523	A	20060809	CN 2004-80019205	20040503
	JP 2006525271	T	20061109	JP 2006-505356	20040503
	AT 373637	T	20071015	AT 2004-730833	20040503
	ES 2293253	T3	20080316	ES 2004-730833	20040503
	US 20070179183	A1	20070802	US 2006-555286	20061017 <--
PRAI	ES 2003-1004	A	20030505		
	WO 2004-EP4670	W	20040503		
OS	MARPAT 143:326627				
GI					



AB The invention relates to phenylalanine derivs. I [G = CO₂H or tetrazolyl; L = a direct bond, NR_c, O, NR_cCO, CONR_c, O₂CNR_c, NR_cCO₂, where R_c = H, alkyl; R₁, R₂ = independently H, (un)substituted (cyclo)alkyl, alkenyl, alkynyl, heterocyclyl, (hetero)aryl, etc.; or NR₁R₂ = (un)substituted heterocyclyl, heteroaryl; R₃, R₄ = H, alkyl; R₅ = (un)substituted (hetero)aryl; R₆ = OH, alkoxy, NO₂, halo, alkylsulfonyl, sulfamoyl, amino, acyl, carboxy, carbamoyl, CN, alkyl, alkenyl, alkynyl, etc.; n = 0-3] and their pharmaceutically-acceptable salts or esters which are α 4 integrin antagonists. For example, reaction of Me (2S)-2-[[[(tert- butoxycarbonyl)amino]sulfonyl]amino]-3-[4-[(2,6- dichlorobenzoyl)amino]phenyl]propionate (preparation given) with benzyl alc. in the presence of PBu₃ and ADDP in THF, followed by saponification with LiOH•H₂O in THF gave (S)-II (43%). In α 4 β 1 adhesion assays, the latter inhibited U-937 cell adhesion to recombinant human soluble VCAM-1 with IC₅₀ values < 100 nM. Thus, I and compns. comprising them are useful for the treatment of inflammatory and immune disorders (no data).

L7 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

AN 2004:996111 CAPLUS Full-text

DN 141:410709

TI Preparation of N-(2-phenylethyl)sulfamide derivatives as integrin α 4 antagonists for treatment of inflammatory and immune disorders

IN Jimenez Mayorga, Juan Miguel; Vidal Gispert, Laura; Warrelow, Graham

PA Almirall Prodesfarma, S.A., Spain

SO PCT Int. Appl., 79 pp.

CODEN: PIXXD2

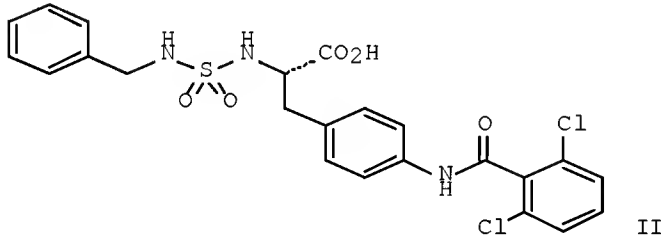
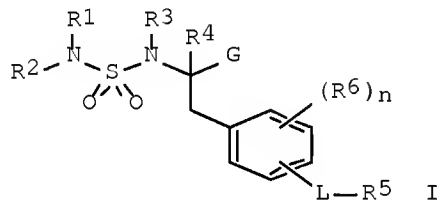
DT Patent

LA English

FAN.CNT 2

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PI	WO 2004099126	A1	20041118	WO 2004-EP4670	20040503
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,				

LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
 NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
 TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
 RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
 AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
 EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
 SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
 SN, TD, TG
 ES 2219177 A1 20041116 ES 2003-1004 20030505
 ES 2219177 B1 20060216
 EP 1622867 A1 20060208 EP 2004-730833 20040503
 EP 1622867 B1 20070919
 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
 IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
 CN 1816523 A 20060809 CN 2004-80019205 20040503
 JP 2006525271 T 20061109 JP 2006-505356 20040503
 US 20070179183 A1 20070802 US 2006-555286 20061017 <--
 PRAI ES 2003-1004 A 20030505
 WO 2004-EP4670 W 20040503
 OS MARPAT 141:410709
 GI



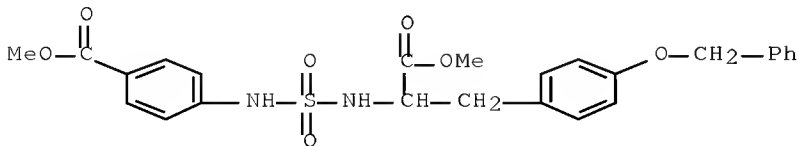
AB Title compds. L-phenylalanine derivs. I [wherein G = CO₂H, tetrazolyl; L = direct bond, NRc, O, NR_cCO, CONRc, OCONRc, NR_cCO₂; Rc = H, alkyl; R₁, R₂ = independently H, (un)substituted (cyclo)alkyl, alkenyl, alkynyl, heterocyclyl, (hetero)aryl, etc.; or NR₁R₂ = (un)substituted heterocyclyl, heteroaryl; R₃, R₄ = H, alkyl; R₅ = (un)substituted (hetero)aryl; R₆ = OH, alkoxy, NO₂, halo, alkylsulfonyl, sulfamoyl, amino, acyl, carboxy, carbamoyl, CN, alkyl, alkenyl, alkynyl, etc.; n = 0-3; and pharmaceutically acceptable salts and esters thereof] were prepared as integrin $\alpha 4$ antagonists. For example, reaction of Me (2S)-2-[[[(tert-butoxycarbonyl)amino]sulfonyl]amino]-3-[4-[(2,6-dichlorobenzoyl)amino]phenyl]propionate (preparation given) with benzyl alc. in the presence of PBu₃ and ADDP in THF, followed by saponification with LiOH•H₂O in THF gave (S)-II (43%). In $\alpha 4\beta 1$ adhesion assays, the latter inhibited U-937 cell adhesion to recombinant human soluble VCAM-1 with IC₅₀

values < 100 nM. Thus, I and compns. comprising them are useful for the treatment of inflammatory and immune disorders (no data).

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d 110 bib abs hitstr

L10 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2008 ACS on STN
AN 2000:41769 CAPLUS Full-text
DN 132:194332
TI Synthesis of 1,2,5-thiadiazolidin-3-one 1,1-dioxide derivatives and evaluation of their affinity for MHC class-II proteins
AU Ducry, Laurent; Reinelt, Stefan; Seiler, Paul; Diederich, Francois; Bolin, David R.; Campbell, Robert M.; Olson, Gary L.
CS Laboratorium fur Organische Chemie der Eidgenossischen Technischen Hochschule, ETH-Zentrum, Zurich, CH-8092, Switz.
SO Helvetica Chimica Acta (1999), 82(12), 2432-2447
CODEN: HCACAV; ISSN: 0018-019X
PB Verlag Helvetica Chimica Acta
DT Journal
LA English
AB 1,2,5-Thiadiazolidin-3-one 1,1-dioxide derivs. were designed by mol. modeling as MHC (major histocompatibility complex) class-II inhibitors. They were prepared from the unsym. N,N'-disubstituted acyclic sulfamides. These N-alkyl-N'-arylsulfamide precursors were synthesized by nucleophilic substitution of either a sulfamoyl chloride or a N-sulfamoyloxazolidinone. Extension of base-induced cyclization methods from aliphatic to aromatic sulfamides gave access to the desired target mols. The N-alkyl-1,2,5-thiadiazolidin-3-one 1,1-dioxide derivs. were also prepared by the oxazolidinone route for coupling to a tetrapeptide fragment. The X-ray crystal structure of 1,2,5-thiadiazolidin-3-one 1,1-dioxide was solved, and the directionality of the H-bond donor (N-H) and acceptor (SO₂) groups of the cyclic scaffold determined. 1,2,5-Thiadiazolidin-3-one 1,1-dioxides were shown to inhibit competition peptide binding to HLA-DR4 mols. in the single-digit millimolar concentration range.
IT 259794-36-0P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of thiadiazolidinone dioxide derivs. and evaluation of their affinity for MHC class-II proteins)
RN 259794-36-0 CAPLUS
CN Tyrosine, N-[[[4-(methoxycarbonyl)phenyl]amino]sulfonyl]-O-(phenylmethyl)-, methyl ester (CA INDEX NAME)



SESSION WILL BE HELD FOR 120 MINUTES

STN INTERNATIONAL SESSION SUSPENDED AT 16:06:13 ON 11 JUL 2008